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S/056/62/043/005/046/058 B125/B104

AUTHORS:

Komarov, L. I., Fisher, I. Z.

TITLE:

On the theory of the Rayleigh scattering of light in fluids

PERIODICAL:

Zhurnal eksperimental noy i teoreticheskoy fiziki, v. 43, no. 5(11), 1962, 1927 - 1933

TEXT: A molecular theory of the intensity and spectrum of the Rayleigh scattering of light in liquids or dense gases is formulated without using the thermodynamical theory of fluctuations. This paper is a transcript of one by L. Van Hove (Phys. Rev., 95, 249, 1954) from the "language" of scattering of neutrons to the "language" of scattering of light. N is the number of molecules contained in the volume V and  $R_i(0)$  (i = 1,2,...,N)

indicates the position of the particles at a certain initial moment,  $R_{\underline{i}}(t)$  is their position at a later moment. The spectral density

 $I'(\mathbf{R}, \omega) = \frac{\alpha^2 \omega^4 N}{2\pi c^4 R^2} I_0 \sin^2 \gamma \int_{(V)} d\mathbf{r} \int_{-\infty}^{\infty} dt \times \exp\left\{i\left(\mathbf{k}_0 - \frac{\omega \mathbf{R}}{cR}\right)\mathbf{r} - i\left(\omega_0 - \omega\right)t\right\} G(|\mathbf{r}|, t),$ (21)

Card 1/3

On the theory of the Rayleigh ...

S/056/62/043/005/046/058 B125/B104

of the intensity of the scattered light is a Fourier transform of a space-time molecular Van Hove function  $G(|\vec{r}|,t)$  of the scattering system.  $\cos \vec{r} = |\vec{E}_0\vec{R}|/R|\vec{E}_0|$ ;  $I_0$  and  $\vec{E}_0$  are the intensity and the electrical vector of the incident radiation.  $G(|\vec{r}|,t)$  is (with an accuracy up to the coefficient  $N^{-1}$ ) the density of the relative probability of finding a certain particle, at the instant t, at a distance  $\vec{r}$  from the initial position of any given particle in the system. After scattering through the argle of  $0^{\circ}$  has been excluded, G has to be replaced by G-1. The integration over  $d\vec{r}$  can be extended over the entire space. Eq. (21) then leads to the formula

$$I'(\mathbf{R},\omega) = \frac{2\pi^2\omega^4N}{c^4R^2} I_0 \sin^2\gamma \int_0^\infty r^2 dr \frac{\sin \varkappa r}{\varkappa r} \int_{-\infty}^\infty dt \left(G(r,t) - 1\right) e^{t\Omega t}. \tag{25}$$

or, with  $G(\vec{r},0) = \delta(\vec{r}) + g(|\vec{r}|)/v$ , to the formula

$$I'(R,\omega) = \frac{\alpha^2 \omega^4 N}{c^4 R^2} I_0 \sin^2 \gamma \delta (\omega - \omega_0) \left\{ 1 + \frac{4\pi}{v} \int_0^\infty (g(r) - 1) \frac{\sin \kappa r}{\kappa r} r^2 dr \right\}. \quad (27).$$

In contrast to the static theory, the present dynamic theory yields a certain definite spectrum of the frequencies in scattering. The formulas Card 2/3

On the theory of the Rayleigh ...

S/056/62/043/005/046/058 B125/B104

(25) and (27) yield equal intensities of the scattered light (aside from corrections of the order of  $(v/c)^2$ ). The polarizability of one molecule in the field of the neighboring molecules is therefore  $\alpha = v^2(-\partial \epsilon/\partial v)_T/4\pi$ , where  $\epsilon$  is the dielectric constant. Up to now, the spectrum of scattered light cannot be calculated from formula (25).

ASSOCIATION: Belorusskiy gosudarstvennyy universitet (Belorussian State University)

SUBMITTED: June 19, 1962

Card 3/3

## "APPROVED FOR RELEASE: 06/13/2000 CIA-RDP86-00513R000413310008-1

S/053/62/076/c03/003/005
B125/B102

AUTHOR: Fisher, I. Z.

TITLE: Present stage of the theory of fluids

PERIODICAL: Uspekhi fizicheskikh nauk, v. 76, no. 3, 1962, 499-518

TEXT: On the basis of papers published from 1935 to 1962 a short survey is given on the present stage of the statistical theory of ordinary fluids with restriction to the classical theory. According to the opinion of the present author, the modern development of this theory was initiated by Yakov Il'ich Frenkel'. There are 1 figure and 46 references:

16 Soviet and 28 non-Soviet.

ZAYTSEVA, A.M.; FISHER, I.Z.

Motion of hydrated ions in solutions. Zhur.strukt.khim. 4 no.1: 3-9 Ja-F '63.

1. Belorusskiy gosudarstvennyy universitet.

(Electrolyte solutions) (Ions-Higration and velocity)

ZATYS	EVA, A.M.; FISHER	, I.Z.		_	
	Effective mass 4 no.2:173-17	of a hydrated ion i	n solution. Zhur stru	kt.khim. (MIRA 16:5)	- 14 - 81 - 81
	1. Belorusskiy	gosudarstvennyy uni IonsMigration and	versitet, Minsk. velocity) (Hydration)		
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ZAYTSEVA, A.M.; FISHER, I.Z.

Mobility of hydrated ions. Zhur.strukt.khim. 4 no.2:261-262
Mr-Ap '63. (MIRA 16:5)

1. Belorusskiy gosudarstvennyy universitet, Minak.
(Hydration) (Ions--Migration and velocity)

	Effect of the hydration of ions on the viscosity of electrolyte solutions. Zhur.strukt.khim. 4 no.3:331-335 My-Je '63.					
	1. Belorusski	y gosudarstvennyy (Electrolyte solu	niversitet, Minsk. tions) (Hydration)	(MIRA 16:	6)	
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#### "APPROVED FOR RELEASE: 06/13/2000 CIA-RDP86-00513R000413310008-1

FISHER, I.Z.; ADAMOVICH, V.I.

Density fluctuations in water. Zhur.strukt.khim. 4 no.6:819-823 N-D '63. (MIRA 17:4)

1. Belorusskiy gosudarstvennyy universitet.

ACCESSION NR: APLOOOL11

5/0046/63/009/004/0427/0433

AUTHORS: Kacharskaya, L. V.; Komarov, L. I.; Fisher, I. Z.

TITLE: Hypersound and slow neutron scattering in liquids

SOURCE: Akusticheskiy zhurnal, v. 9, no. 4, 1963, 427-433

TOPIC TAGS: hypersound neutron wave diffraction, liquid hypersound neutron wave diffraction, high frequency hypersound, hypersound neutron scattering, slow neutron scattering, hypersound slow neutron scattering, hypersound neutron scattering spectrum, hypersonic radiation, neutron scattering, neutron, scattering, hypersound

ABSTRACT: The conditions leading to neutron wave diffraction on hypersounds in liquids are analyzed, and conditions for building an experimental apparatus specified. Slow neutron scattering is found to be the most suitable because of the requirements of very high frequencies for the determination of hypersounds in fluids. The expression for the neutron energy E and momentum p during and after the scattering is represented by

 $E - E_0 = \pm u(\Omega) |p - p_0|,$ 

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ACCESSION NR: APL	0070070450	imentally determin			
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ASSOCIATION: Belor University)	usakiy gosmdaratvenn	ymy universitet, M	insk (Byelorussier	State	
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ADAMOVICH, V.I.; KUNTSEVICH, I.M.; FISHER, I.Z.

Short-range order near an excited molecule in a liquid. Zhur.
fiz. khim. 37 no.11:2568-2570 N\*63. (MIRA 17:2)

5/056/63/044/002/025/065 B102/B186

/UTHOR:

Fisher,

TITLE:

Localized excitons in fluids

FERIODICAL:

Zhurnal eksperimental noy i teoreticheskoy fiziki, v. 44;

no. 2, 1963, 541-547

TEXT: The author develops a hydrodynamic theory for the motion of a localized exciton, considered as a quasiparticle, in a fluid. The localized exciton, which in the microscopic theory is considered as a specifically ordered region in the neighborhood of an electron-excited molecule moving along the exciton, is regarded in the hydrodynamic theory as a small dense region in the fluid. The only assumption made is the necessity for such conditions that a localized exciton is not only produced but also conserved for a sufficiently long period. The above-mentioned dense region is characterized by  $\varrho(\vec{r}) = \varrho_0 \exp\left[-\vec{r}(\vec{r})/kT\right]$  where  $\varrho_0$  is the normal density

and Y(r) is the mean selfconsistent force potential, considered as given, and  $\vec{Y}(\vec{r}) = \vec{Y}(|\vec{r}|)$ . The force acting around the center of the exciton is Card 1/4

Localized excitons in fluids

S/056/63/044/002/025/065 B102/B186:

given by  $\vec{K}(\vec{r}) = \frac{1}{V} \nabla p = -\frac{c^2}{kT} \nabla \vec{\Psi}(|\vec{r}|)$  (per mass unit);  $p_r$  is the pressure and  $c^2 = (\partial p/\partial q)_T$ . For an ideal fluid

$$\rho(\mathbf{r}) = \rho_0 \exp\left\{-\frac{\overline{\Psi}(|\mathbf{r}|)}{kT} + \sigma(\mathbf{r})\right\}$$
 (3)

$$K(\mathbf{r}) = -\frac{c^4}{kT} \nabla \widetilde{\Psi}(|\mathbf{r}|) + c^2 \nabla \sigma(\mathbf{r}).$$
 (4).

Since the kinetic energy can be set up as  $E_{\rm kin} = \frac{1}{2} \int \varrho(\vec{v} - \vec{U})^2 dv \approx \frac{1}{2} \varrho_0 U^2 (\vec{v}_W)^2 dv$ , the factor multiplied by  $U^2/2$  can be taken as the effective mass. The motion is considered in a system in which the exciton center is at rest, i.e. the fluid has a stationary flow with the velocity  $\vec{U}$  at infinity;  $\vec{V} = \vec{U} + U \nabla \psi(\vec{r})$ ,  $\Delta \psi = \frac{1}{kT} \vec{Y}(r) \cos \theta$ ,  $\vec{V}(r) = \begin{cases} \mathcal{E}(r/R-1) & r < R \\ 0 & r > R \end{cases}$ . One obtains  $M_{\rm eff} = \frac{2\pi}{15} \left(\frac{\mathcal{E}}{kT}\right)^2 \varrho_0 R^3$ , or, if the total mass in the sphere of radius R is  $M = 4\pi \varrho_0 R^3/3$ ,  $M_{\rm eff} = \frac{1}{30} \left(\mathcal{E}/kT\right)^2 M \ll M$ . Subsequently the exciton motion is investigated for a viscous fluid. The characteristic length Card 2/4

8/056/63/044/002/025/065
Localized excitons in fluids B102/B186

L =  $\frac{U}{v_0c^2} (\frac{4}{3}\eta + \frac{4}{5})$  takes displacive ( $\eta$ ) and volume ( $\frac{4}{5}$ ) viscosity into account.

$$\varphi(x,y,z)=\sum_{n=0}^{\infty}(-L)^n\frac{\partial^n\varphi_0}{\partial z^n}.$$
 (24)

will be a solution of the kinetic equation

$$\Delta \left( \varphi + L \frac{\partial \varphi}{\partial z} \right) = \frac{1}{RT} \frac{\partial \overline{\Psi}}{\partial z}, \qquad (20).$$

For small but finite L,  $\psi(x,y,z)=\psi_0(x,y,z-L)$ . When energy discipation due to viscosity is taken into account; the exciton is subjected to the frictional force  $\hat{F}_{fr} = -h(-\vec{v})$ , where  $(-\vec{v})$  is the velocity of an exciton whose coefficient of friction is

$$h = \int \left[ 2\eta \left( \frac{\partial^2 \varphi}{\partial x_l \partial x_k} \right)^2 + \left( \zeta - \frac{2}{3} \eta \right) (\Delta \varphi)^2 \right] dV. \tag{27}.$$

Card 3/4

Localized excitons in fluids

\$/056/63/044/002/025/065 B102/B186

For slow motions: it

$$h = \frac{4\pi}{9} \left(\frac{59}{60} \eta + \zeta\right) \left(\frac{\varepsilon}{kT}\right)^2 R. \tag{29},$$

which, compared with the Stokes law  $h=6\pi\gamma\,R^*$ , yields the inequality  $R^*\leqslant R$ ; R is the effective radius of an exciton considered as a hard sphere and R is the linear dimension of the region occupied by the exciton. The relaxation time in the case of translational motion can be estimated from  $\tau = M_{ff}/h$ , and the diffusion coefficient from  $D = kT/h = kT\tau/M_{eff}$ .

ASSOCIATION:

Belorusskiy gosudarstvennyy universitet (Belorussian State

University)

SUBMITTED:

June 26, 1962 (initially) October 22, 1962 (after revision)

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ACCESSION NR: AP4040373

S/0185/64/009/004/0349/0354

AUTHOR: Komarov, L. I.; Fisher, I. Z.

TITLE: Neutron and Optical Spectra as Sources of Information on the Motion of Molecules in Liquids /Paper presented at the Shestoye Soveshchaniye po Fizike Zhidkogo Sostoyaniya Veshchestva, Sixth Conference on the Physics of the Liquid.

SOURCE: Ukrayins'ky\*y fizy\*chny\*y zhurnal, v. 9, no. 4, 1964, 349-354

TOPIC TAGS: liquid state, liquid state physics, liquid molecular motion, molecular radial density distribution, molecular correlation function, many-body problem, Van Hove time molecular distribution function, Rayleigh scattering, neutron scattering, fluctuation theory

TRANSLATION: The Van Hove time molecular distribution formalism is introduced and applied to the description of such kinetic properties of condensed phases as correlation and autocorrelation functions and fluctuations. The possibilities of obtaining information about these functions from experimental data are studied

Card 1/2

ACCESSION NR: AP4040373

in detail. It is shown that the Rayleigh scattering spectrum can be used to determine the asymptotic behavior of the Van Hove functions with respect to distance and time. Low energy neutron scattering by liquids is considered in detail with a view toward extracting information about nolecular distributions. The high frequency limit of sound propagation in liquids is considered, and the possibility of the existence of transverse waves in liquids is likewise treated. The phonon approach of solid state usage is modified to treat the neutron scattering problem in liquids. Though neutron scattering differential cross-sections and the angular distribution of Rayleigh scattering are not expected to be precisely conformal, they are expected to be closely related. Very low energy neutron scattering is recommended as a tool for studying the nature of molecular behavior which gives rise to the dispersion of elastic and kinetic coefficients in liquids at the high frequency limit. Orig. art. has 17 numbered equations.

ASSOCIATION: Belorusskiy Gosuniversitet, Minsk (Byelorussian State University)

SUBMITTED:

SUB CODE: OP, GP DATE ACQ: 13May64

NO REF SOV: 006

ENCL: 00

OTHER: 001

Card 2/2

ACCESSION NR: AP4040375

\$/0185/64/009/004/0379/0383

AUTHOR: Fisher, I. Z.

TITLE: On the Theory of Diffusion Near the Critical Point Paper presented at the Shestoye Soveshchaniye po Fizike Zhidkogo Sostoyaniya Veshchestva, Sixth Conference on the Physics of the Liquid State of Matter, Kiev, 1963.

SOURCE: Ukrayins'ky\*y fizy\*chny\*y zhurnal, v. 9, no. 4, 1964, 379-383

TOPIC TAGS: Thermodynamics, diffusion, liquid diffusion, critical point, diffusion coefficient, critical point diffusion, Chaager kinetic coefficient, Brownian motion, liquid kinetics, entropy

TRANSLATION: A treatment, more exact than those extant, is presented for the theory of diffusion processes in a binary-system near the critical point. The diffusion coefficient of any component is proportional to the partial derivative of its chemical potential with respect to its concentration:

$$D (c) \sim \left(\frac{\partial \mu}{\partial c}\right)_{T}$$

(1)

Card 1/3

ACCESSION NR: AP4040375

At the critical point, however

$$D(c) \sim \left(\frac{\partial u}{\partial c}\right)_{T} = 0.$$

Thus, diffusion should cease at the critical point. Any supposition based on the absence of molecular mixing, even for a single state of a substance, would contradict a molecular-kinetic approach. This difficulty justifies the investigation, which is based on the deviation of the entropy from its equilibrium value. The expression for entropy contains as one of its terms the gradient of the off-equilibrium concentration deviation, Sc. Differential equations are obtained for "critical diffusion", and Green's functions of these equations are constructed for one- and three-dimensional cases. If Sc is treated as a "phase", its "center of gravity" changes with a characteristic time

(21)

according to the present treatment for the critical point. The usual diffusion

Card 2/3

ACCESSION NR: AP4040375

 $\langle |x| \rangle = \text{const} \sqrt{2Dt}$ 

(22)

where t is the time and D H. Thus, the slow nature — slow, yet still non-zero — of the diffusion is explained by the small value of H as well as the th dependence. Also in contrast to the usual diffusion theory, a fundamental difference is seen to exist between macroscopic diffusion and Brownian motion of individual particles: the Green's functions have negative values at particulat transition probabilities for Brownian particles. Orig. art. has: 35 formulas.

ASSOCIATION: Belorusskiy Gosuniversitet, Minsk (Byelorussian State University)

SUBMITTED: 00

DATE ACQ: 13May64

ENCL: 00

SUB CODE: JP

NO REF SOV: 003

OTHER: 000

Card 3/3

## "APPROVED FOR RELEASE: 06/13/2000 CIA-RDP86-00513R000413310008-1

PICHER, I.Z.; SOSHINA, H.V.; ZAYTSEVA, A.M.

Density aderofluctuations in simple liquids. Ukr. fiz. zhur. 9 no.5: 476-480 My '64. (NIRA 17:9)

1. Belorusskiy gosudarstvennyy universitet, Minsk.

Comments on the kinetic theory of liquids. Zhur, fiz. khim.
38 no.3:778-780 Mr '64. (MIRA 17:7)

1. Belorusekly gosudarstvennyy universitet.

FISHER I.Z.; ZAYTSEVA, A.M.

Effect of the hydration of ions on the volume viscosity of electrolyte solutions. Dokl. AN SSSR 154 no.5:1175-1178 F'64.

1. Belorusskiy gosudarstvennyy universitet im. V.I.Lenina.

Predstavleno akademikom I.I. Chernyayevym.

## KOMAROV, L.I.; FISHER, I.Z.

TO THE PROPERTY OF THE PROPERT

Neutron and optical spectra as sources of information on the motion of molecules in liquids. Ukr. fiz. zhur. 9 no.4:349-354 Ap 164. (MIRA 17:8)

1. Belorusskiy gosudarstvennyy universitet, Minsk.

FISHER, I.Z.

Theory of diffusion near the critical point. Ukr. fiz. zhur. 9 no.4:379-393 Ap 64. (MIRA 17:8)

1. Belorusskiy gosudarstvennyy universitet, Minsk.

ZAYTSEVA, A.M.; FISHER, I.Z.

Hydration theory. Ukr. fiz. zhur. 9 no.4:383-387 Ap '64.

(MIRA 17:8)

1. Belorusskiy gosudarstvennyy universitet, Minsk.

FISHER. I.Z.

Laminar nature of a diffusion flow near the critical point in a binary solution. Dokl. AN SSSR 158 no.5:1075-1077 0 64.

(MIRA 17:10)

1. Odesskiy gosudarstvennyv universitet im. I.I.Mechnikova. Predstavleno akademikom M.A.Leontovichem.

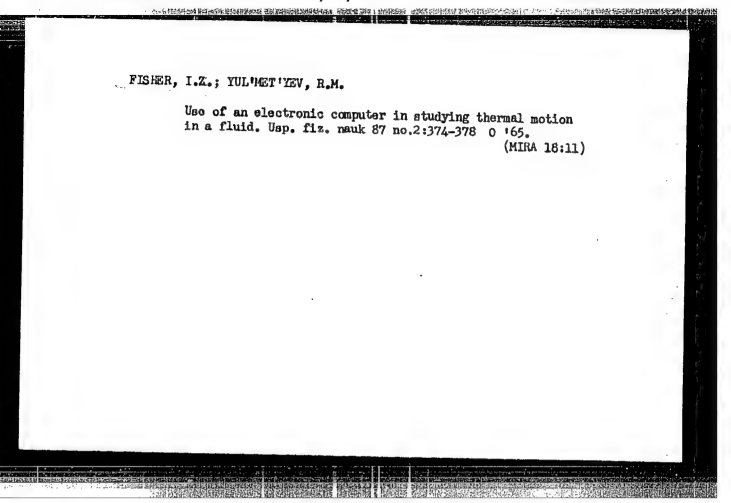
FISHER, I.Z.; KOVALENKO, N.P.

Effect of walls on the fluctuation level near the critical point, Zhur.fiz.khim. 39 no.10:2569-2571 0 165.

(MIRA 18:12)

1. Odesskiy gosudarstvennyy universitet imeni Mechnikova. Submitted June 6, 1964.

## "APPROVED FOR RELEASE: 06/13/2000 CIA-RDP86-00513R000413310008-1



FISHER, K,

FISHER, K.; SKARICA, R. Electric transmission linesiin Croatia. p. 249

THE PROPERTY OF THE PROPERTY O

Vol. 9, no. 4/5, Apr./May 1956 ELEKTROPRIVERDA TECHNOLOGY Beograd

Sor East European Accession, Vol. 6, no.3, March, 1957

APPROVED FOR RELEASE: 06/13/2000 CIA-RDP86-00513R000413310008-1"

## "APPROVED FOR RELEASE: 06/13/2000 CIA-RDP86-00513R000413310008-1

PARENKO, P., FISHER, L., SAMARIN, A., RUDENKO, P.

This is most important in educational work. Sov. profsoiuzy
17 no.18:21 S '61.

(Community centers)
(1rade unions)

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DEKHTYAR, B.; FISHER, L.; UDATOV, A. (g.Mogocha, Chitinskoy obl.);
TOLETIY, P. (g. Yagotin, Kiyeyakaya obl.); SOLODOVNIKOV, I.
(Primorskiy kray); MUN'KO.T. (g. Zaporozh'ye)

Letters and correspondence. Sov.profsoiuzy 17 no.22:42-44 N
'61.

1. Spetsial'nyy korrespondent zhurnala "Sovetskiye profsoyusy?.
(Community centers, Mobile)
(Ural Mountain region—Callisthen'cs)
(Adult education)
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NAZARENKO, P. (Astrakhanskaya oblast'); KIL'DIBEKOV, V. (g.Slobodskoy, Kirovskaya oblast'); DEVYATOVSKIY, M. (g.Orsk); SERGIYENYA, K. (g.Khar'kov); FISHER, L.; BELYAYEV, A.; VENGEROV, A.; KRAVTSOV, S. (g.Khar'kov)

Readers relate, advise and criticise. Sov. profsoiuzy 18 no.15:26-28 Ag '62. (MIRA 15:7)

1. Neshtatnyy korrespondent zhurnala "Sovetskiye profsoyuzy" (for Nazarenko, Sergiyenya, Vengerov). 2. Sotrudnik gorodskoy gazety "Leninskiy put'" (for Kil'dibekov). 3. Sotrudnik neshtatnogo otdela oblostnogo kimiteta profsoyuza rabochikh metall rgicheskoy promyshelnnosti (for Devyatovskiy). 4. Predsedatel' Ermiteta profsoyuza elektromekhanicheskogo zavoda, g.Khar'kov (Yor Kravtsov). (Socialist competition) (Ust'-Kamenogorsk—Housing) (Kharkov—Electric equipment industry)

## "APPROVED FOR RELEASE: 06/13/2000 CIA-RDP86-00513R000413310008-1

FISHER, L. (Krasnaya Polyana, Moskovskaya obl.)

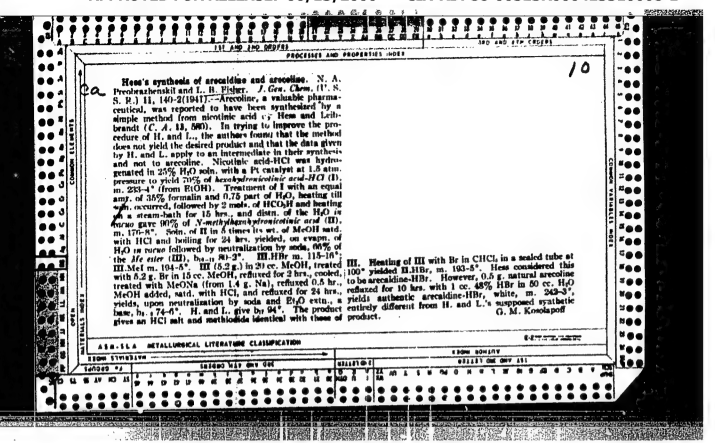
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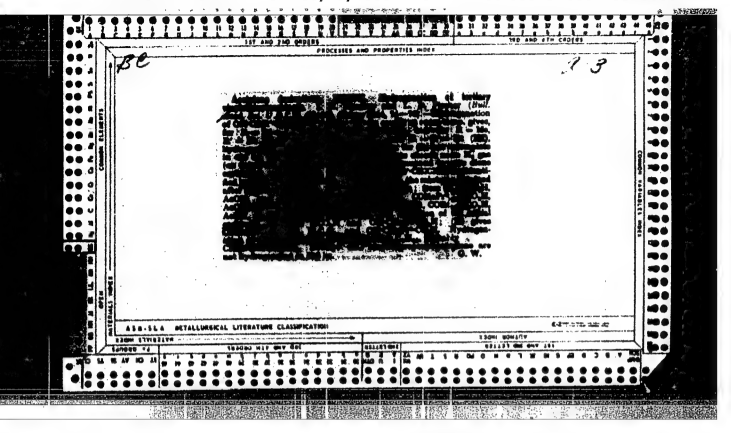
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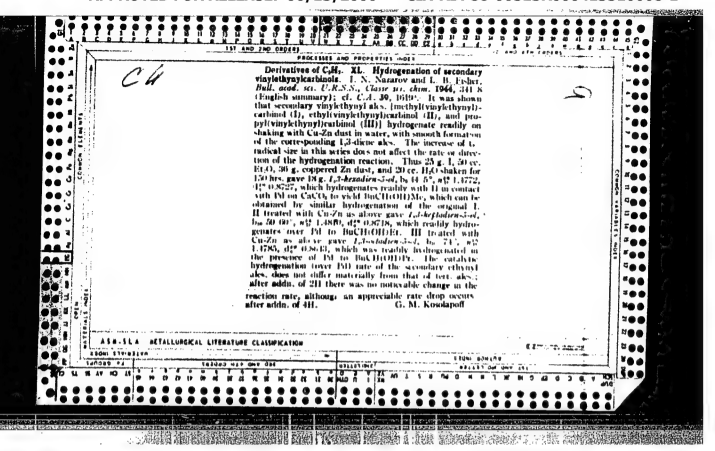
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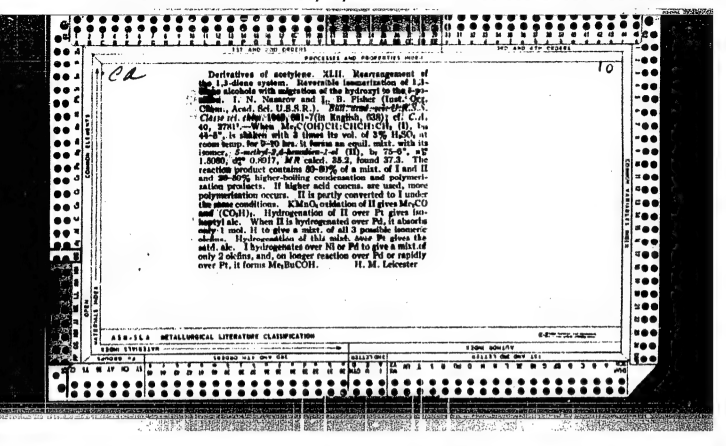
(Krasnaya Polyana—Gotton spinning) (Socialist competition)

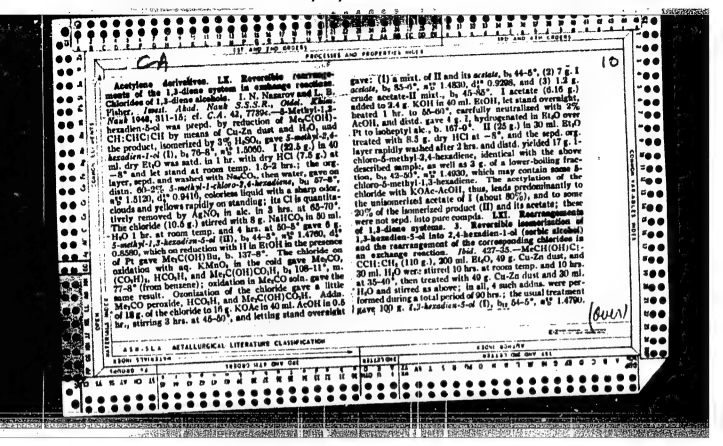
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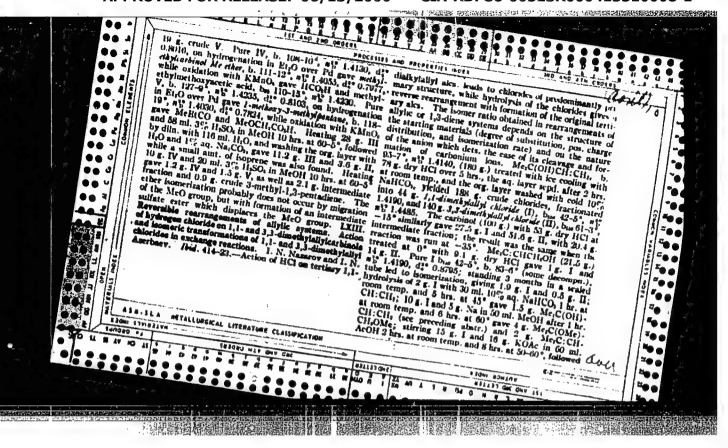












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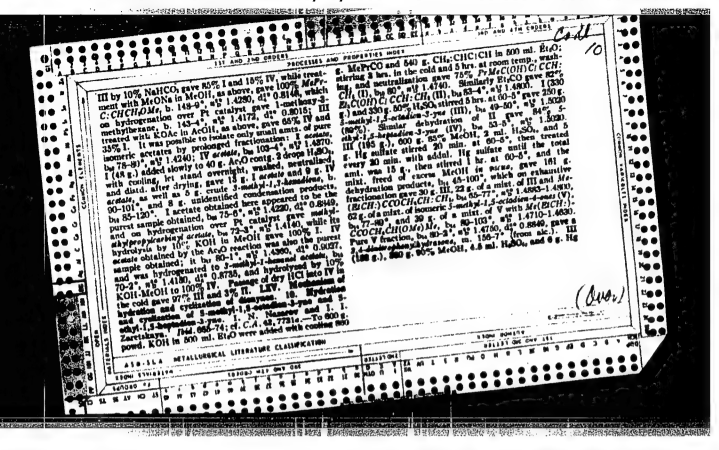
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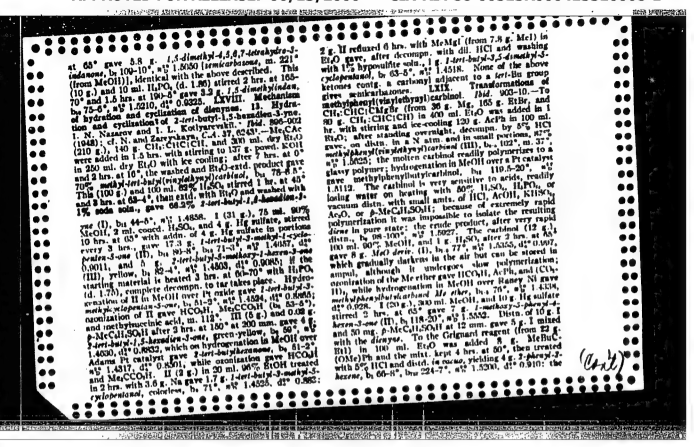
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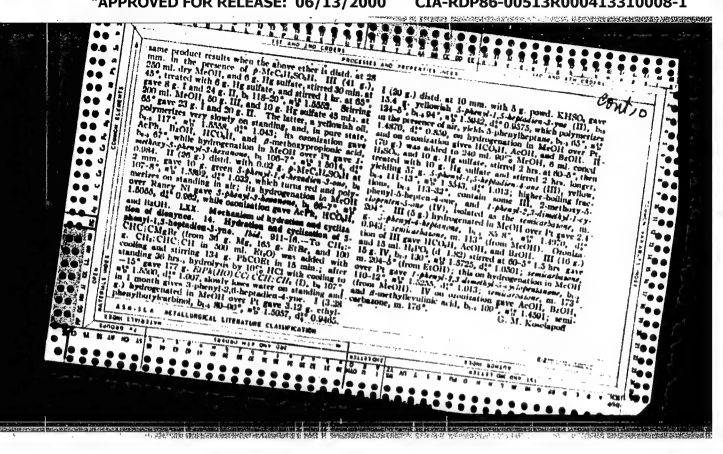
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(d. 1.78) was used under similar conditions (the semicar-land) of the cyclopentenous gave HCO<sub>H</sub> and ittle methylicensine and sold of the cyclopentenous gave HCO<sub>H</sub> and ittle methylicensine and sold of the cyclopentenous gave HCO<sub>H</sub> and ittle methylicensine acid (isolated as the RCO<sub>H</sub>), and a little methylicensine acid (isolated as the RCO<sub>H</sub>), and a little methylicensine acid (isolated as the RCO<sub>H</sub>), and a little methylicensine acid (isolated as the RCO<sub>H</sub>), and a little methylicensine acid (isolated as the RCO<sub>H</sub>), and a little methylicensine acid (isolated as the RCO<sub>H</sub>), and a little methylicensine acid (isolated as the RCO<sub>H</sub>), and a little methylicensine acid (isolated as the RCO<sub>H</sub>), and a little methylicensine acid (isolated as the RCO<sub>H</sub>), and a little methylicensine acid (isolated as the RCO<sub>H</sub>), and a little methylicensine acid (isolated as the RCO<sub>H</sub>), and a little methylicensine acid (isolated as the RCO<sub>H</sub>), and a little methylicensine acid (isolated as the RCO<sub>H</sub>), and a little methylicensine acid (isolated as the RCO<sub>H</sub>), and a little methylicensine acid (isolated as the RCO<sub>H</sub>), and a little methylicensine acid (isolated as the RCO<sub>H</sub>), and a little methylicensine acid (isolated as the RCO<sub>H</sub>), and a little methylicensine acid (isolated as the RCO<sub>H</sub>), and a little from the unother liquor), is that of the IX isomer observation as a little from the unother liquor), is that of the IX isomer observation as also yielded 10 g. of the other isomer of IX, by 08-0°, as also yielded 10 g. of the other isomer of IX, by 08-0°, as also yielded 10 g. of the other isomer of IX, by 08-0°, as also yielded 10 g. of the other isomer of IX, by 08-0°, as also yielded 10 g. of the other isomer of IX, by 08-0°, as also yielded 10 g. of the other isomer of IX, by 08-0°, as also yielded 10 g. of the other isomer of IX, by 08-0°, as also yielded 10 g. of the other isomer of IX, by 08-0°, as also yielded 10 g. of 08-0°, as also yielded 10 g. of 08-0°, as also yielded 10 g. of 08-0°, as yielded 10 g. of 08-0°, as yield sulfate stirred 2 lus. at 00°, then treated in 2 portions with 10°, lig sulfate with a 2-br. interval, stirred 6 hrs. at 00°, and worked up as above, gave 212°, products, predominantly the above Mol hidone (VI), but 86°, at 1.4540, 40°, 0.9243. V (7.8°, fraction but 80-2°) in 20° mi. BtOH (0.9243. V (7.8°, fraction but 80-2°) in 20° mi. BtOH (0.9243. V (7.8°, fraction but 80-2°) in 20° mi. BtOH (0.9243. V (7.8°, fraction but 80-2°) in 20° mi. BtOH (0.9243. V (2.8°, fraction but 80-2°) in 20° mi. BtOH (0.9243. V (2.8°, fraction but 80°, 5°, mich yi-4-octament, to general states of the grave 3 g. of previously to give a cryst. axime or semicarhanone. Ononination of to give a cryst. axime or semicarhanone. Ononination of the seminar results were obtained by cyclication with HCl at 60°-5° or with p-McCalt-SO-H at 120-20°. V (9°,) and 10°, 12°, in 10°, in .. .. -.. .. •• .. .. .. .. .. .. 









FISHER, L. B.

USSR/Chemistry - Acetylene, Derivatives Chemistry - Isomerization Jul-Aug 48

"Acetylene Derivatives, No 61," I. N. Nasarov, L. B. Fisher, Inst Org Chem, Acad Sci USSR, 8½ pp

"Iz Ak Nauk, SSSR Otdel Khim Nauk" No 4

Investigates reversible isomerisation of 1, 3-hexadiene-5-ol into 2,4-hexadiene-1-ol (sorbic alcohol) under the influence of dilute sulfuric acid. Investigates reaction of hydrogen chloride with these two diene alcohols and also hydrolysis and acetylisation of the chlorides thus obtained. Describes reversible regrouping of 1,3-diene system observed in the exchang reactions mentioned. Submitted 7 Jul 1947

PA 8/49T17

FISHER. L. B.

USSR/Chemistry - Acetylehe, Derivitives Chemistry - Hydrogenation

Jul/Aug 48

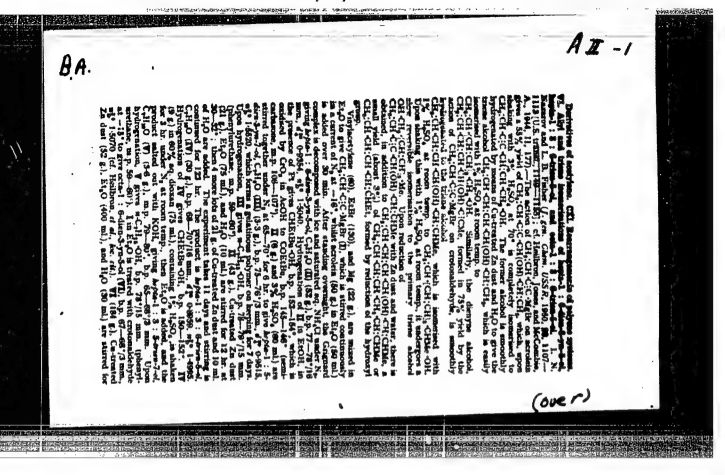
"Acetylene Derivatives, No 81," I. N. Nazarov, L.B. Fisher, Inst Org Chem, Acad Sci, USSR, 84 pp

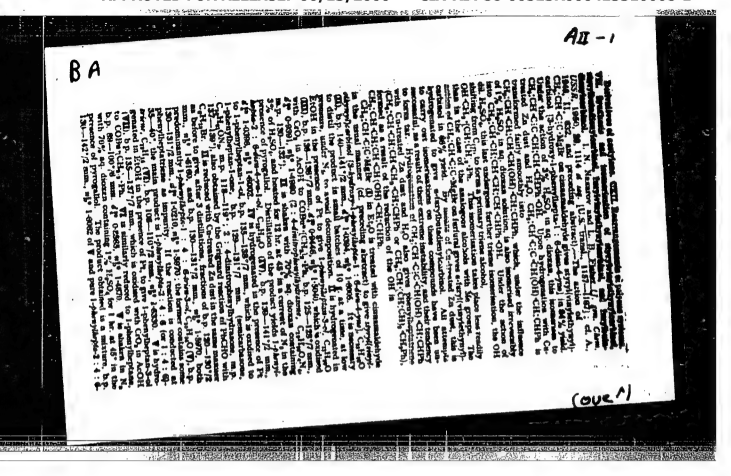
一次的设计 对他的自己的指数**对比较级对**相通过第一位的 第二种人类的现在分词形式,这种关键,但是是不是是一种,不是不是一个

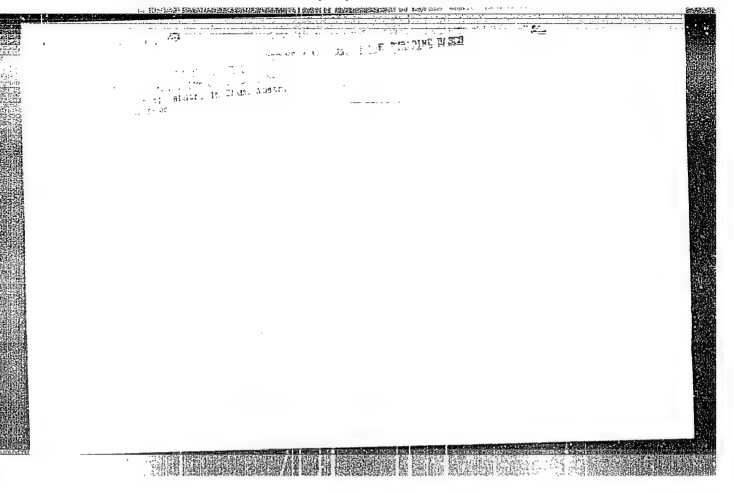
"Iz Ak Nauk SSSR, Otdel Khim Nauk" No 4.

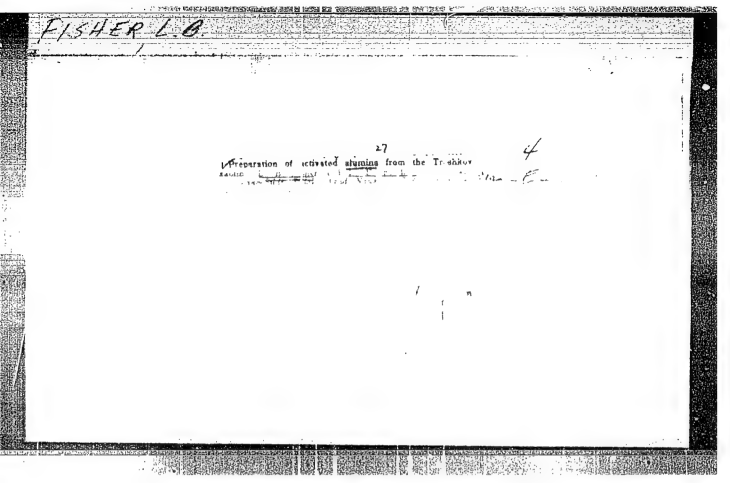
Describes hydrogenation of phenyl-vinylethylene-carbinol (I) into 5-phenyl, 3-pentadiene-5-ol (II) by means of copperized zinc dust and water. Achieved irreversible isomerisation of 5-phenyll-1, 3-pentadiene-5-ol into 5-phenyl-2, 4 pentadiene-1-ol (III) by the action of diluted sulfuric scid. Investigated action of hydrogen chloride on diene alcohols II and III, and also the hydrolysis and esterification of the primary chloride thus obtained. Submitted 6 Jun 1947.

PA 8/49T18









PLYAREVSKIY, I.L.; FISHER, L.B.  Hydrogenation of gaseous olefins under pressure, Zhur. prikl.			
khim. 29 no.11:1	im. 29 no.11:1756-1757 N '56. (MIRA 10:6)		
l. Laboratoriya organicheskogo sintesa Vostochno-Sibirskogo filiala Akademii nauk SSSR. (Hydrogenation) (Olefine)			birskogo

FISHER, L.B.; BELLEN'KATA, A.P.; MARUSHKIN, M.N. [deceased],

Catalytic dehydrogenation of butane to butene over catalysts obtained from local raw materials. Isv. vost. fil. AM SSER no.1153-59 157.

1. Vostochno-Sibirskiy filial AM SSER,

(Butane) (Behydrogenation) (Kaolin)

FISHER, L.B.: TERPENOVA, M.P.; KOTLYAREVSKIY, I.L.

Gatalytic dehydrogenation of 2,3-dimethylbutane, Izv. vost. fil.
AN SSSR no.9:53-56 '57. (MIRA 11:1)

1. Vostochno-Sibirskiy filial AN SSSR,
(Butane) (Dehydrogenation)

AUTHOR: Fisher, L. B. (Irkutsk)

14 27-5-5/6

TITLE:

Amines of the Acetylene Series (Aminy atset:lenovogo

ryada)

PERIODICAL:

Uspekhi Khimlis 1958; Vol. 27, Nr. 5, pp. 589-621 (USSR)

ABSTRACT:

In the introduction the author points out that beside the methods of synthesis of acerylene-amines methods based upon Mannich's reaction (reaktsiya Mannikha) recently gained importance. In the years before World War Two a great number of processes was worked out which are based upon the reaction of acetylene with ammonta and with amines. Later the works in this field were intensively continued. Then in the present paper the methods are discussed in individual sections: the catalytic methods, the method of the synthesis of acetylene-amines (according to Rappa; references 12 - 15), further the synthesis of aminobutynes published in 1949. A special section is devoted to Mannich's reaction. Among others it is pointed out that Marshak and his collaborators in the investigation of the condisions of the synthesis of tertiary acetylene-amines by means of Mannich's reaction found that an excess of formaldehyde must be obtained at the end of the

Card 1/3

Amines of the Acetylene Series

74-07-3-3/6

reaction. Further data on the mechanism of the reaction of acetylene compounds with amines and formeldehydes follow. The works by Dornow and Isha (reference 43), Marszak (references 44, 19, 41, 45, 46), Navarov and Shvekbgeymer (reference 54) are discussed. A special section deals with the problem of the production of acetylene samines by means of substitution reactions and so talled mixed methods. The works by Ferrit and Weart (Teferance 66) Mandach and Kulkes (reference 69), as well as Perveyer (reference 78, 79) are referred to. The next section deals with the properties and the use of sostylene smines. The sythor discusses the production of a number of disubstituted vertiary acetylene. amines by the condensation of sodium substituted diethylaminopropyne with different haloud alkyls, whereby a number of dimethylaminoalkanes could be synthesized (Parsell and Pollard, reference 8t). The author also describes the synthesis of primary, secondary, and tertiary amino-slochols by means of the interaction or monosubstituted amintacetylene compounds with organic exides alsochydes and ketones in the presence of acdium amide in liquid ammoura. Then follows an

Card 2/3

APPROVED FOR RELEASE: 06/13/2000 CIA-RDP86-00513R000413310008-1"

Amines of the Acetylene Series

74-27-5-3/6

enumeration of further syntheses: The production of new acetylene-1,4-aminoalcohols, of acetylene-aminoesters and other compounds. The isomerization of N-substituted compounds of 2-aminobutyne-3 to N-substituted 2-aminobutadiene-1,3 derivatives; the hydrogenation of acetylene and its derivatives is described in detail. The 24 tables added to the paper offer a good survey. There are 24 tables, 108 references, 25 of which are Soviet.

1. Acetylene derivatives--Synthesis

contraction of the second seco

Card 3/3

PISHER, L.B.; TERPUGOVA, M.P.; KOTLYAREVSKIT, I.L.

Dehydrogenation of butane di-derivatives; studying the first stage of 2,3-dimethylbutane dehydrogenation. Inv.Sib.otd. AW SSSR no.9:32-38 '58. (MIRA 11:11)

1. Vostochno-Sibirekiy filial Akademii nauk SSSR. (Butane) (Dehydrogenation)

5(1,3) SOV/153-2-4-25/32 AUTHORS: Kotlyarevskiy, I. L., Fisher, L. B., Zanira, A. S., Terpugova, M. P., Volkov, A. N., Shvartsberg, M. S. TITLE: Synthesis of Several Moncmers on Alumochromium Catalysts PERIODICAL: Izvestiya vysshikh uchebrykh zavedeniy. Khimiya i khimicheskaya tekhnologiya, 1959, Vol 2, Nr 4, pp 608 - 613 (USSR) ABSTRACT: A report on this paper was given at the All-Union Conference on "Ways of Synthesis of Initial Products for the Production of High Polymers" which took place in Yaroslavl' from September 29 to October 2, 1958. The results of the catalytic synthesis of 2,3-dimethyl-butadiene-1,3 as well as of monomers of the type of p-xylene from acetylene derivatives are given. The substance mentioned at first can be used in the production of special rubber types (Ref 1). It could not be obtained by the production methods so far used. The diagram (see Diagram) being worked at by the authors consists of two stages. The first one (Ref 2) yielded not more than 15% of the end product. In spite of numerous patents (Ref 3), a thorough description of reaction conditions is still missing. Therefore, the authors determined Card 1/4 the optimum conditions of isobutane alkylation with ethylene

Synthesis of Several Monomers on Alumochromium Catalysts SOV/153-2-4-26/32

in the presence of aluminum chloride. Under these conditions  $(48 - 51^{\circ}, 7 - 9 \text{ atm}, 1.5 \text{ kg isobutane}, 145 \text{ g ethylene}, 40 \text{ g})$ CoH5Cl, 5 & AlCl3, 2 hours) the yield of 2,3-dimethyl-butane rises to 47%. The reaction is very sensitive to temperature (only 1/3 of the alkylate yield at  $35^{\circ}$ ). The catalyst can be used 5 to 6 times without reducing the quantity of alkylate or of 2,3-dimethyl-butane. Ethylene has to be added during the whole process: its partial pressure must not exceed 0.5 - 0.7 atm, or otherwise the 2,3-dimethyl-butane content in the alkylate decreases rapidly. The addition of 1-3%  $C_2H_5Cl$  accelerates the process. Isobutane alkylation with ethylene at a high temperature proceeds at a pressure of 100 - 200 atm according to a radical mechanism, and is accelerated by radical donors. It was carried out by the authors in the presence of C2H5Cl (fcr the first time) in a special, continuously working plant. The alkylate yield (computed for ethylene) reached 170-180% under optimum conditions (450°, 200 atm, reaction time 25 minutes, weight ratio isobutane : ethylene = 10 : 1, initiator quantity 2.5%). 2,2-Dimethyl-butane (32-35% of the alkylate) was the

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Synthesis of Several Monomers on Alumochromium Catalysts SOV/153-2-4-26/32

main product: 2,3-dimethyl-butane amounted to 8-10% of the alkylate. The preparation of 2,3-dimethyl-butadiene-1,3 from 2,3-dimethyl-butane proceeds in two stages: a) Dehydration with the formation of two olefines: 1) tetramethyl-ethylene, and 2) 2.3-dimethyl-butene-1. b) Dehydration of the olefines to the end product. This reaction has not been considered in publications (except for the reference in the patent (Ref 5)). The authors investigated the first stage thoroughly on an alumochromium-potassium catalyst. The specifications for its production were provided by M. N. Marushkin (deceased) (IOKh AN SSSR = Institute of Organic Chemistry AS USSR). This catalyst showed the highest activity at 550°. The catalyzate reached a 87% yield with a content of unsaturated hydrocarbons of 60-67%. The investigation of the second stage has not yet been completed. An aromatization diagram of divinyl-acetylene-hydrocarbons is given. Several mono-, bi-, and tricyclic hydrocarbons with a prescribed structure were produced on the basis of the diagram. A diagram of the reactions of the synthesis of p-xylene from acetylene and acetone is suggested. Since there is no demand for acetone in the USA because of increasing phenol production,

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Synthesis of Several Monomers on Alumochronium Catalysts S07/153-2-4-26/32

and its price according to the Sergeyev method is already 1/3 - 1/4 of the present one, the synthesis method mentioned above might become useful for industry. In conclusion, a simple way of synthesis for polyphenyl systems is suggested. Moreover, the Favorskiy reaction is mentioned in the paper. There are 11 references, 5 of which are Sovict.

ASSOCIATION: Vostochno-Sibirskiy filial SO AN SEER (East Siberian Branch of the Siberian Department of the Academy of Sciences, JECR)

Card 4/4

KOTLYAREVSKIY, I.L.; VOLKOV, A.N.; FISHER, L.B.

Laboratory method for producing 2,3-dimethylbutane by the alkylation of isobutane by ethylene. Izv. Sib. otd. AN SSSR no.3:62-66 '59. (MIRA 12:8)

1. Vostochno-Sibirskiy filial Sibirskogo otdeleniya Akademii nauk SSSR.

(Butane) (Ethylene) (Alkylation)

KOTLYAREVSKIY, I.L.; VOLKOV, A.N.; FISHER, L.B.

Alkylation of butane and isobutane by ethylene. Izv.Sib.otd.
AN SSSR no.4:64-70 59. (HIRA 12:10)

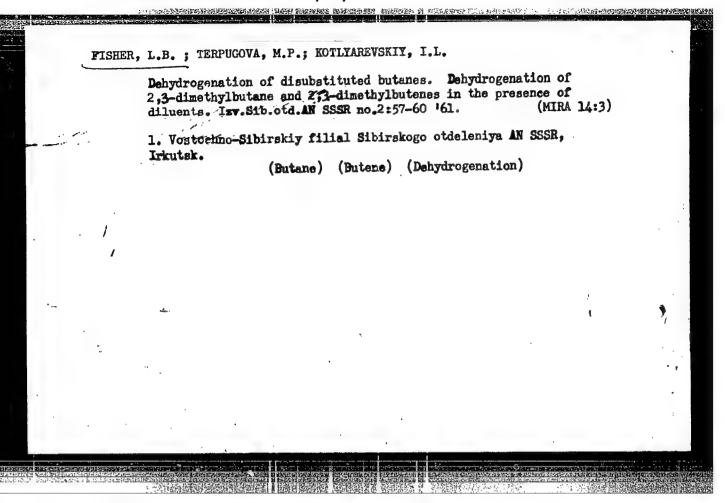
1. Vostochno-Sibirskiy filial Sibirskogo otdeleniya Akademii nauk SSSR. (Butane) (Ethylene) (Alkylation)

KOTLYAREVSKIY, I.L.; FISHER, L.B.; ZANIMA, A.S.; TERPUGOVA, M.P.;
VOLKOV, A.M.; SHVARTSBERG, M.S.

Synthesis of some monomers over aluminochronium catalysts.

Isv.vys.ucheb.mav.; khim.i khim.tekh 2 no.4:608-613
159. (MIRA 13:2)

1. Vostochno-Sibirskiy filial Srednessiatskogo otdeleniya
AN SSSR. (Butadiene) (Catalysis)



CIA-RDP86-00513R000413310008-1

20270 S/062/61/000/010/015/018 B106/B101

15.8063

Kotlyarevskiy, I. L., Fisher, L. B., and Domnina, Ye. S.

AUTHORS: TITLE:

Oxidative polycondensation of diacetylene compounds

PERIODICAL:

Akademiya nauk SSSR. Izvestiya. Otdeleniye khimicheskikh

nauk, no. 10, 1961, 1905 - 1907

TEXT: In a previous paper (Ref. 1: I. L. Kotlyarevskiy, L. B. Fisher, A. A. Dulov, A. A. Slinki, Izv. AN SSSR. Otd. khim. n. 1960, 950) the example of p-diethynvl benzene (I) showed that oligomers with conjugate aromatic and acetylene bonds can be obtained by oxidative condensation to a linear diacetylene compounds. After this paper, another one describing a similar synthesis was published (Ref. 2: see below). The physical properties of the poly-p-diethynyl benzene thus synthesized are similar to properties of the poly-p-diethynyl benzene thus synthesized are similar to those of semiconductors and ferromagnetics. In the present paper, the examination of this reaction is continued. The corresponding polymers for which, in analogy to II, the structures VII, VIII, IX, and X are suggested, were obtained by oxidative polycondensation of 1,4-diethynyl-1,4-dihydroxy-cyclohexane (III), 9,10-diethynyl-9,10-dihydroxy-9,10-dihydroxy-Card 1/6

### "APPROVED FOR RELEASE: 06/13/2000

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Oxidative polycondensation of ...

20270 S/062/61/000/010/015/018 B106/B101

anthracene (IV), 9,10-diethynyl-9,10-dihydroxy-9,10-dihydro phenanthrene (V), and 9,10-diethynyl anthracene (VI).

X

Card 2/6

Oxidative polycondensation of ...

S/062/61/000/010/015/018 B106/B101

The initial p-diethynyl benzene (I) was obtained by two methods: (1) from the dehydrogenation products of p-diethyl benzene (Ref. 4: A. A. Balandin, N. I. Shuykin, G. M. Marukyan, I. I. Brusov, R. G. Seymovich, G. K. Lavrovskaya, V. K. Mikhaylovskiy, Zh. prikl. khimii, 32, No. 11; 2566, 1959), which were brominated in carbon tetrachloride at -12 - -15°C. The resulting tetrabromide was filtered off, washed with carbon tetrachloride, and recrystallized from chloroform (melting point: 156 - 157°C). Hydrogen bromide was then split off from the above tetrabromide by a method of R. Deluchat (Ref. 7: Ann. chimie 11, 1, 181 (1934)), and p-diethynyl benzene (melting point: 95°C) was thus obtained. 2) From p-diacetyl benzene by catalytic hydrogenation to 1,4-di(-1-hydroxy-ethyl)-benzene, by dehydration to p-divinyl benzene, and by bromination and further treatment of the latter as described under 1). The monomers III - V were obtained by condensation of acetylene and diketones in liquid ammonia (Ref. 5: W. Ried, H. J. Schmidt, Chem. Ber. 90, 2499 (1957)). Compound VI was obtained by aromatization of IV (Ref. 6: W. Ried, H. J. Schmidt, A. Urschel, Chem. Ber. 91, 2472 (1958)). Cuprous chloride was used as a catalyst for polycondensation, and aqueous dioxane, aqueous alcohol, and pyridine (Ref. 2: see below) were used as solvents. Card 4/6

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25278 S/062/61/000/010/015/018 B106/B101

Oxidative polycondensation of ...

solution of the catalyst and of the corresponding monomer in the solvent used was shaken together with oxygen at room temperature until gas absorption came to an end. The reaction mixture was decomposed with water, and the deposited product was washed with water until the chloride was removed. Oligomer II is orange-red (at n = 6) or bright-yellow (at n = 5), and insoluble in water and conventional organic solvents. Oligomer VII is bright-colored, polymers VIII and IX are darkish brown powders, and polymer X is black. These polymers are all insoluble in conventional organic solvents at room temperature. Compounds VIII and IX contain traces of copper but no chlorine, and polymer X contains chlorine. There are 8 references: 3 Soviet and 5 non-Soviet. The two references to English-language publications read as follows: Ref. 2:
A. S. Hay, J. Organ. Chem. 25, 1275 (1960); C. A. Hochwalt, Pat. USA 2390368 (1954); Chem. Abstrs, 1878 (1946).

ASSOCIATION: Institut khimii Sibirskogo otdeleniya Akademii nauk SSSR (Institute of Chemistry of the Siberian Department of the

Academy of Sciences USSR)

Card 5/6

33374 5/190/62/004/002/002/021 B110/B101

15.8340

AUTHORS:

Kotlyarevskiy, I. L., Fisher, L. B., Dulov, A. A.,

Slinkin, A. A., Rubinshteyn, A. M.

TITLE:

Synthesis and some physical properties of poly-p-diethynyl

benzene

PERIODICAL:

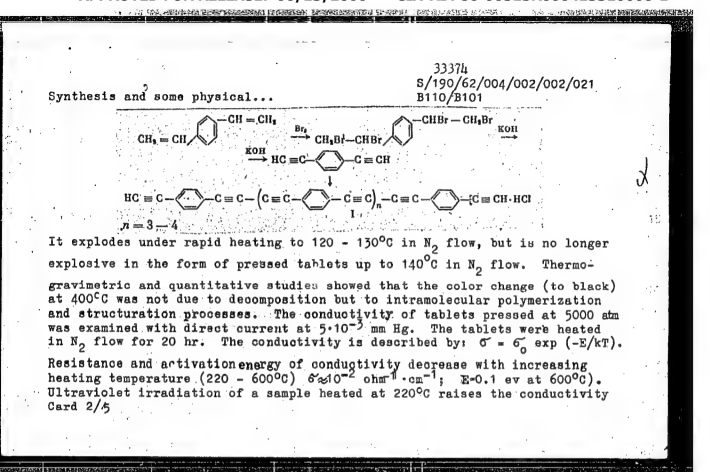
Vysokomolekulyarnyye soyedineniya, v. 4, no. 2, 1962,

174 - 181.

TEXT: Poly-p-diethinyl benzene with alternating ternary bonds and phenylene rings was synthesized from p-diethinyl benzene according to Yu. S. Zalikind (Zh. obshch. khimii, 6, 530, 1936). The diethyl benzene mixture obtained during styrene production was dehydrogenated to divinyl benzene, brominated, dehydrobrominated, and polycondensed in water-alcohol or water-dioxane at 20 - 40°C in the presence of CuCl, NH<sub>4</sub>Cl, and 0<sub>2</sub> to orange-red, powdery

oligomer (I) insoluble in water and organics:

Card 1/3



33374

Synthesis and some physical...

S/190/62/004/002/002/021 B110/B101

reversibly by some orders. This effect decreases with increasing heating temperature (400°C) and disappears at 500 and 600°C completely. The sign of the thermo-emf and the reversible resistance decrease during oxygen adsorption confirm the hole character of the conductivity. The specific magnetic susceptibility of the initial oligomer determined between 20 and 160°C at H = 3500 - 4500 cersteds was  $\chi = -0.4 \cdot 10^{-6}$ , after pressing at 5000atm:  $\chi = -0.2 \cdot 10^{-6}$ . The maximum number of unpaired electrons exists on heating to 220°C, maximum  $\chi$  value at 400°C, while ferromagnetic H dependence on  $\chi$ was observed. The intensities of the epr signal as dependent on heating (2 hrs) in vacuo and N<sub>2</sub> (0.5% O<sub>2</sub>) pass through a maximum at  $\sim$  220°C. X-ray studies with an YP(-55(URS-55) device showed increasing crystal formation (favored by pressing) with increasing heating temperature. The electric and magnetic properties of slightly heated amorphous samples are determined by individual unpaired electrons and energetic barriers between the loosely bound, conjugated sections while ultraviolet irradiation increases the number of current carriers. At higher temperatures, the individual conjugated sections are combined to microcrystalline domains, and the number of electrons which have not yet entered the domain of strong interaction Card 3/5

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Synthesis and some physical...

decreases. At a certain stage, further crystal growth gives rise to formation of diamagnetic graphite structures. At 400, 500, 600°C, electric resistance and activation energy of conductivity decrease with increasing heating temperature due to the presence of strongly interacting electrons. Two types of structurally different polymers are likely to exist. The conversion of the orange-colored, explosive initial polymer at 200°C is likely to proceed according to:

$$C \equiv C - \left(C \equiv C - \left(C \equiv C\right)_{n} - C \equiv C\right) - C \equiv C - \left(C \equiv C - \left(C \equiv C\right)_{n} - C \equiv C\right) - C \equiv C - C \equiv$$

Card 4/5

33374

Synthesis and some physical...

5/190/62/004/002/002/021 B110/B101

At higher temperatures, domains are formed with ferromagnetic electron interaction due to cross linking which are superposed by diamagnetic interaction on further crystallization. Tal'roze is mentioned. There are 4 figures, 4 tables, and 14 references: 11 Soviet and 3 non-Soviet. The two references to English-language publications read as follows: A. S. Hay, J. Org. Chem., 25, 1275, 1960; D. D. Eley et al., Disc. Faraday Soc., 28, 55, 1959.

ASSOCIATION:

Institut khimii Vostochno-sibirskogo filiala AN SSSR (Institute of Chemistry of the East Siberian Branch AS USSR). Institut organicheskoy khimii im. N. D. Zelinskogo AN SSSR (Institute of Organic Chemistry imeni N. D. Zelinskiy AS USSR)

SUBMITTED:

January 30, 1961

Card 5/5

KHAGANOV, Ye.I., otv. red.; FISHER, L.B., red.; POPOV, K.V., red.; STENNIKOV, M.G., tekhn. red.

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[Materials for the conference of young scientists; on the tenth anniversary of the East-Siberian Branch of the Academy of Sciences of the U.S.S.R.] Materialy k konferentsii molodykh nauchnykh sotrudnikov; k 10-letiiu Vostochno-Sibirskogo filiala AN SSSR. Blagoveshchensk. No.3. [Chemistry and metallurgy] Khimiia i metallurgiia. 1960. 93 p. (MIRA 17:2)

1. Akademiya nauk SSSR. Vostochno-Sibirskiy filial, Irkutsk.

EFA(5) 0/EWT(m)/EPF(c)/FPR/EWP(j)/T Pc-4/Pr-4/Ps-4/Pt-10 s/0062/64/c00/006/1090/1095 FUTE IN VP AP4041169 AUDICE: Fisher, L. B.; Kotlyarevskiy, I. L.; Domning. Ye. S.; Trotsenko, Z. B. MITTLE: Highly unsaturated polymers. Communication 9. Synthesis and polycondensation of discetylenic derivatives of phenanthrene BOURCE: AN SSSR. Izv. Seriya khimicheskaya, no. 6, 1964, 1090-1095 FOFIC TAGS: discetylenic polymer, discetylenic phenenthrene derivative, synthesis, polycondensation, unsaturated polymer, acetylenic oligomer, conjugated way winer polymer, conjugated polymer, organic semiconductor, semiconducting polymer ABSTRACT: As a continuation of research to determine the effects of monomer on the physical properties of oligoners, oligoners were synthesized in a gation was retained or interrupted, and copolymers were synthesized in saving intermipted conjugation alternated with various ted units. on many . - ), lo-dietyty 1-3, 10-dinydrophener tarens . . . was and for m Cord 1/2

ACCESSION NR: AP4041169

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and powdered KOE; 9,10-diethynylphenanthrene (II) was made by aromatizing III.

There subjected to oxidative polycondensation in pyridine with catalyst

there subjected to oxidative polycondensation in pyridine with catalyst

there was intuited. Copolymers of I and II. and of I and it enterprise

between 'IV), and of III and IV were similarly made. The IR spectra and EH data

for the products are tabulated. "Klectron paramagnetic resonance spectra were

chtained in the laboratory by Y. Y. Yovnevolsko, V. K. Yernolsyev, H. M. Gazhin

hapovalov, whom the authors thank." Originant, has a takes and 2

sets of equations.

ARRYTAPING. Institut khimicheskoy kinetiki i goreniya Sibirakogo otdeleniya

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Card 2/2

FISHER, L.B.; KOTLYAREVSKIY, I.I.; DOMNINA, Ye.S.; TROTSENKO, Y.F.

。 [1] \$ 14.15 \$16.15 \$

Highly unsaturated polymers. Report No.9: Synthesis and polycondensation of diacetylene derivatives of phenanthrene. Izv. AN SSSR. Ser. khim. no.6:1090-1095 Je 164.

(MIRA 17:11)

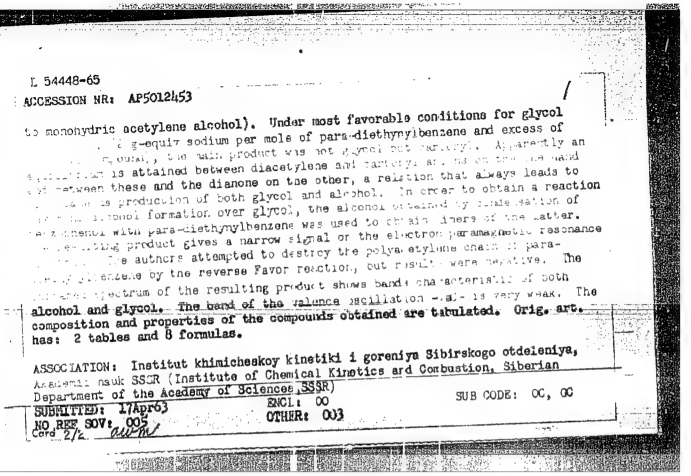
1. Institut khimicheskoy kinetiki i goreniya Sitirskogo otdeleniya AN SSSR.

FISHER, L.B.; KOTLYAREVSKIY, I.L.; ANDRIYEVSKAYA, E.K.

Mannich reaction with p-diethynylbenzene derivatives. Izv. AN SSSR. Ser. khim. no.8:1543-1545 Ag '64. (MIRA 17:9)

1. Institut khimicheskoy kinetiki i goreniya Sibirskogo otdeleniya AN SSSR.

ACCESSION NR: AP5012453  BYTHORS: Fisher, L. B.; Kotlyarevskiy, I. L.  TITLE: Highly unsaturated polymers. Communication 11. Memohydric and dihydric alcohols derived from para-diethynylbenzene  SOURCE: AN SSSR. Izvestiya. Seriya khimicheskaya, no. L, 1965, 692-697  TOPIC TAOS: unsaturated compound, alcohol, benzol, contensation reaction, organic synthesis, IR spectrum  ABSTRACT: The behavior of paradiethynylbenzens in the Newland reaction was studied. Since acetylene reacts with sodium and sodium anide in liquid ammonia to form a memoration derivative that will not react further with sodium (or Nakii;) because the compound of the Nakii and reaction parmits production of the reactylene alcohols a vithout always that it is acetylene if are tylene alcohols a vithout always that it is acetylene alcohols a vithout always that it is acetylene alcohols a vithout always that are action, the authors undertook condensation of the compound with acetone, cyclohexanone, and benzophenone. In all cases a principal product (70-80%) was glycol (in addition	ACCESSION NR: AP5012453  UR/0062/65/000/004/0692/0697 5h7.362+5h1.6  Z7  AUTHORS: Fisher, L. B.; Kotlyarevskiy, I. L.  TITLE: Highly unsaturated polymers. Communication 11. Memohydric and dihydric alcohols derived from para-diethynylbenzene  SOURCE: AN SSSR. Izvestiya. Seriya khimicheskaya, no. b, 1965, 692-697  TOPIC TAGS: unsaturated compound, alcohol, benzol, contensation reaction, organic synthesis, IR spectrum  IBSTRACT: The behavior of paradiethynylbenzens in the Newland reaction was studied in liquid ammonia to form a since acetylene reacts with sodium and sodium amide in liquid ammonia to form a serium derivative that will not react further with sodium (or Nahii) because the compound of the content of the compound with acetion permits production of the compound with acetions, the investigate the behavior of para-diethynylbenzene in the Newland reaction, the investigate the behavior of para-diethynylbenzene in the Newland reaction, the			75.23
ATTHORS: Fisher, L. B.; Kotlyarevskiy, I. L.  TITLE: Highly unsaturated polymers. Communication 11. Memohydric and dihydric alcohols derived from para-diethynylbenzene (  SOURCE: AN SSSR. Izvestiya. Seriya khimicheskaya, no. k, 1965, 692-697  TOPIC TAGS: unsaturated compound, alcohol, benzol, comiensation reaction, organic synthesis, IR spectrum  ABSTRACT: The behavior of paradiethynylbenzene in the Mewland reaction was studied Since acetylene reacts with sodium and sodium anide in liquid ammonia to form a memoration derivative that will not react further with sodium (or NaNitz) because the compound are action, the Newland reaction parmits production of the reactly sylbenzene in the Newland reaction, the investigate the behavior of para-diethynylbenzene in the Newland reaction, the investigate the behavior of para-diethynylbenzene in the Newland reaction, the investigate the behavior of para-diethynylbenzene in the Newland reaction, the	ANTHORS: Fisher, L. B.; Kotlyarevskiy, I. L.  TITLE: Highly unsaturated polymers. Communication 11. Memohydric and dihydric alcohols derived from para-diethynylbenzene  SOURCE: AN SSSR. Izvestiya. Seriya khimicheskaya, no. h, 1965, 692-697  TOPIC TAGS: unsaturated compound, alcohol, benzol, condensation reaction, organic synthesis, IR spectrum  ABSTRACT: The behavior of paradiethynylbenzens in the Newland reaction was studied since acetylene reacts with sodium and sodium amide in liquid ammonia to form a since acetylene reacts with sodium and sodium amide in liquid ammonia to form a series with sodium and sodium and entering the secure of		UR/0062/65/000/004/0692/0697	
TITLE: Highly unsaturated polymers. Communication 11. Monohydric and dihydric alcohols derived from para-diethynylbenzene  SOURCE: AN SSSR. Izvestiya. Seriya khimicheskaya, no. 4, 1965, 692-697  TOPIC TAGS: unsaturated compound, alcohol, benzol, commensation reaction, organic synthesis, IR spectrum  ABSTRACT: The behavior of paradiethynylbenzene in the Mewland reaction was studied. Since acetylene reacts with sodium and sodium amide in liquid ammonia to form a series acetylene reacts with sodium and sodium amide in liquid ammonia to form a series acetylene saturation, the Newland reaction permits production of acetylene alcohols without always the production of liquid ammonia to form a series acetylene alcohols without always the production of liquid ammonia to form a series acetylene alcohols without always the production of liquid ammonia to form a series acetylene alcohols without always the series acetylene in liquid ammonia to form a series acetylene alcohols without always the series acetylene alcohols without always the series acetylene, and series acetyl	TITLE: Highly unsaturated polymers. Communication 11. Memohydric and dihydric alcohols derived from para-disthynylbenzene  SOURCE: AN SSSR. Izvestiya. Seriya khimicheskaya, no. 4, 1965, 692-697  TOPIC TACS: unsaturated compound, alcohol, benzol, contensation reaction, organic synthesis, IR spectrum  ABSTRACT: The behavior of paradiethynylbenzene in the Hewland reaction was studied. Since acetylene reacts with sodium and sodium amide in liquid ammonia to form a memoration derivative that will not react further with sodium (or Nakhiz) because the least of the least of paradiethynylbenzene paradiethynylbenzene in the Newland reaction, the least of derivatives in the least reaction, the authors undertook condensation of the compound with acetone, cyclohexanone, and benzophenone. In all cases a principal product (70-8%) was glycol (in addition		26	
SOURCE: AN SSSR. Isvestiya. Seriya khimicheskaya, no. 1, 1965, 692-697  TOPIC TAGS: unsaturated compound, alcohol, benzel, contensation reaction, organic synthesis, IR spectrum  ABSTRACT: The behavior of paradiethynylbenzene in the Newland reaction was studied. Since acetylene reacts with sodium and sodium anide in liquid ammonia to form a newsodium derivative that will not react further with sodium (or Nakhi) because newsodium derivative that will not react further with sodium (or Nakhi) because in the newland reaction permits production of the reaction, the Newland reaction, the liquid ammonia to form a nacetylene alcohols without always allowed the production of the compound with acetone, cyclohexanone, and	TOPIC TAGS: unsaturated compound, alcohol, benzol, contensation reaction, organic synthesis, IR spectrum  ABSTRACT: The behavior of paradiethynylbenzens in the Newland reaction was studied.  Gince acetylene reacts with sodium and sodium anide in liquid ammonia to form a newsodium derivative that will not react further with sodium (or Nakhi) because the implete saturation, the Newland reaction permits production of the reactylene alcohols without always and state and givens.  In vertigate the behavior of para-diethynylbenzene in the Newland reaction, the authors undertook condensation of the compound with acetone, cyclohexanone, and benzophenone. In all cases a principal product (70-80%) was glycol (in addition	TITLE: Highly unsaturated polymer	rs. Communication 11. Memohydric and dihydric	
ABSTRACT: The behavior of paradiethynylbenzens in the Newland reaction was studied.  Since acetylene reacts with sodium and sodium amide in liquid ammonia to form a monascium derivative that will not react further with sodium (or NaWi) because monascium derivative that will not react further with sodium (or NaWi) because monascium derivative saturation, the Newland reaction permits production of macetylene alcohols without always with the section, the monascium the behavior of para-diethynylbenzene in the Newland reaction, the monascium of the common with acetone, cyclohexanone, and	ABSTRACT: The behavior of paradiethynylhenzens in the Newland reaction was studied.  Gince acetylene reacts with sodium and sodium amide in liquid ammonia to form a new sectium derivative that will not react further with sodium (or Nakii2) because the interpolate saturation, the Newland reaction permits production of the compound with acetylene alcohols without always that is the gradient in the investigate the behavior of para-diethynylbenzene in the Newland reaction, the authors undertook condensation of the compound with acetone, cyclonexamone, and benzophenone. In all cases a principal product (70-80%) was glycol (in addition	A		
Since acetylene reacts with sodium and sodium and the liquid amount of liquid amount of the companies of the section parmits production of the reactylene alcohols without always the liquid and gradual acetylene alcohols of derivatives. The liquid and gradual in the section, the liquid acetylene	Since acetylene reacts with sodium and sodium and the little of the ambient and the product of the little of the l	TOPIC TAGS: unsaturated compound, synthesis, IR spectrum	, alcohol, benzol, contenuation reaction, organic	
	Card 1/2	Since acetylene reacts with sodium nonrectium derivative that will no new prime acetylene all interestly, yielding two in resettigate the behavior of pa	on and sodium and of it in the data and sodium (or Nakki2) because on, the Newland reaction parmits production of looks a without always aiming the land gayous, are distinguished in the Newland reaction, the fithe company with acetone, cyclohexanone, and	



EWT(1)/EWT(m)/EPA(w)-2/T/EWP(t)/EWP(b)/ENA(m)-2/EWA(c) IJP(c) UR/0386/65/001/005/0054/0057 AP5016286 ACCESSION NR: 50 Fisher. L. M. 4 Sharvin, Yu. V.; AUTHOR: Observation of focused electron beams in a metal SOURCE: Zhurnal eksperimental noy i tekhnicheskoy fiziki. Pis'ma v redaktsiyu. TITLE: Prilozheniye, v. 1, no. 5, 1965, 54-57 TOPIC TAGS: electron beam, electron motion, electron spectrum ABSTRACT: The authors have performed the experiment proposed by one of them earlier (Sharvin, ZhETF v. 48, 984, 1965), aimed at producing and observing in a metal with large mean free path electron beams starting from a definite point of the sample and focused by longitudinal magnetic field in another point of the sample. A disgram of the experiment is shown in Fig. 1 of the Enclosure. The experiment was carried out at 2K with two thin points of tin wire (60  $\mu$  in diameter) were soldered to a single crystal plate of high purity. The current was 200 ma. The sample was placed in a magnetic field which could be varied in magnitude and direction. In the absence of a magnetic field, the measured voltage was quite small because of the large conductivity of the sample. When the magnetic field was turned on, a signal appeared in the circuit, and its magnitude increased when the field became Card 1/3

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perpendicular to the sample. The signal increased approximately in proportion to the square of the magnetic field, but in addition exhibited periodically repeated maxima, which could be attributed to the focusing of definite groups of electrons, accelerated by the electric field. The magnitude of the effect is in satisfactory agreement with the estimates made in earlier papers. Upon focusing, the electrons execute an integral number of revolutions, (moving along helical paths) as they move beneath the ends of the metal. The relation between the observed peaks and the Fermi surface of the metal is briefly discussed. "We thank P. L. Kapitsa for interest in the work." Orig. art. has: 2 figures.

ASSOCIATION: Institut fizicheskikh problem im. S. I. Vavilova Akademii nauk SSSR

(Institute of Physics Problems, Academy of Sciences, SSSR)

SUBMITTED: 29Apr65

ENCL: 01

SUB CODE: NP. MM

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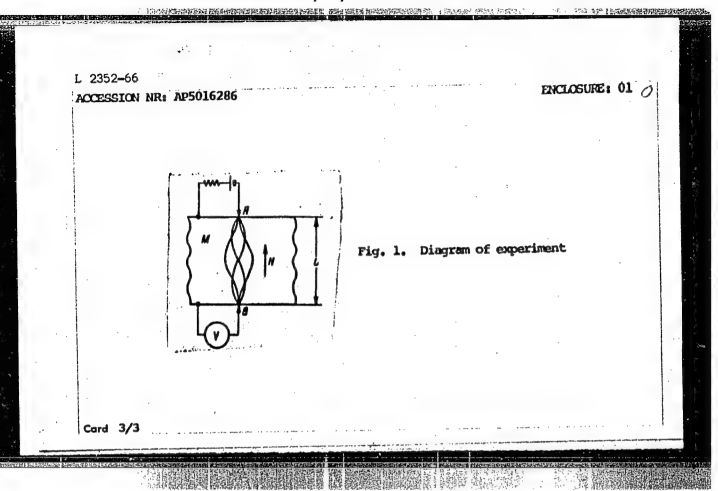


Figure 1. Stylesveriv, I.L.

Righly unsaturated polymers. Report No.11; Monoatomic and diatomic alcoholo, derivatives of p-diethynylbanzene. Izv.

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AN SSSR.

FishER L.I.

## KUNIVASKAIA S. B., FISHER L.I.

Proba Nesterova pri nektorykh zabolevanijakh vmutrennikh organov.
/Hesterova test in certain disenses of internal organy/
arkh. 23:2 Har-Apr 51 p. 63-7.

1. Of the Propedeutic Therapeutic Clinic (Director--Prof. I. I. Tavetkov), Saratov Medical Institute. CLML Vol. 20, No. 10 Oct 1951

## FISHER, L. I.

Cutaneous vascular tests in hypertension. Klin. med., Noskva 29 no.7:32-35 July 1951. (CIML 21:1)

1. Docent. 2. Of the Faculty Therapeutic Clinic (Director Prof. L. A. Varshamov), Saratov Medical Institute.

FISHER, L.I.

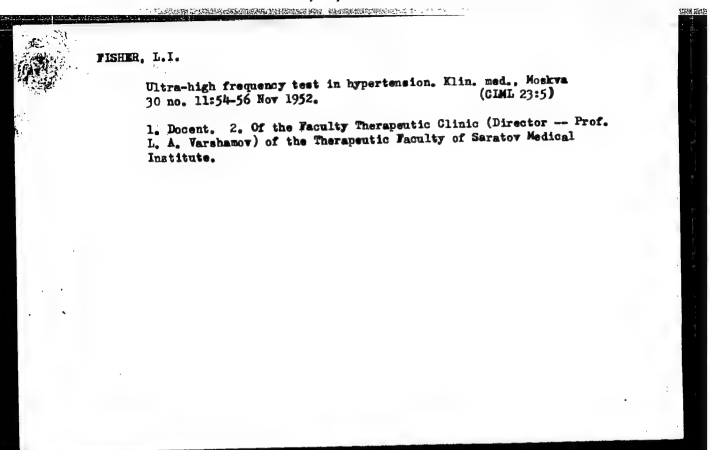
Electrical sensitivity of the eye in hypertension. Klin. med., Hoskva 30 no.3:83 Mar 1952. (CLML 22:2)

1. Docent. 2. Of the Faculty Therapeutic Clinic (Director -- Prof. L. A. Varshamov), Saratov Medical Institute.

VARSHAMOV, L.A.; FISHER, L.I.; IL'IGHEVA, N.S.

Sleep therapy in gastric and duodenal ulcer. Klin. med., Moskva 30 no.9: 65-70 Sept 1952. (GLML 23:2)

1. Professor for Varshamov; Docent for Fisher. 2. Of the Faculty Therapeutic Clinic (Director -- Prof. L. A. Varshamov), Saratov Medical Institute.



FISHER, L.I., dotsent; VARSHAMOV, L.A., professor.

Types of neural reactions and clinical characteristics of hypertension. Terap.arkh.25 no.4:37-48 Jl-Ag '53. (MLRA 7:2)

1. Iz fakul'tetskoy terapevticheskoy kliniki lechebnogo fakul'teta (direktor - professor L.A. Varshamov) Saratovskogo meditsinskogo instituta. (Hypertension) (Nervous system)

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FISHER, L.I., dotsent (Saratov); VARSHAMOV, L.A., professor, direktor.

Autonomic changes in sleep therapy of hypertension. Klin.med. 31 no.9:67-70 S '53.

1. Fakul'tetskaya terapevticheskaya klinika lechebnogo fakul'teta Saratovskogo meditsinskogo instituta. (Hypertension) (Nervous system) (Sleep)

FISHER, L.I., dotsent (Saratov)

Use of physiotherapeutic factors in protective inhibition therapy.

Klin.med. 32 no.9:41-45 5 '54. (MLRA 7:12)

1. Is fakul'tetskoy terapevticheskoy kliniki (dir. prof. L.A.Varshamov) lechebnogo fakul'teta Saratovskogo meditsinskogo instituta.

(SIMEP, therapeutic use)

FISHER JL I

USSR/Human and Animal Physiology - Blood Circulation.

R-5

Abs Jour

: Rererat Zhur - Biologiya, No 16, 1957, 70750

Author

: L.I. Fisher.

Title

: The Effect of "UVCh" on the Functional State of the

Central Nervous System in Hypertensia.

Orig Pub

: Vopr. kurortol. fizioterapii i lecheb. fiz. kul'tury,

1956 7/2, 35-41

Abstract

: In 50 hypertensive patients, the functional state of Central Nervous System was studied before and after the effect of the action of "UVCh" pole on the region of the soles for a period of 15-20 minutes. Verbal experiment, the determination of the electrical sensitivity of the eye and of optical chronaxy, the acuteness of feeling and pain sensitivity, motor chronaxymetry, measurement of skin conductivity, sensory chronasymetry, and plethysmography indicated that in therapy with "UVCh" the inhibitory processes in the Central Nervous System are intensi-

fied.

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Faculty Therapeuties Clinic, Hospital Jacuity, Sacator Med. Inst.

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FISHER, L.I., dotsent

Chronaximetric indexes in hypertension and changes following sleep therapy. Terap.arkh. 28 no.2:33-40 '56. (MLRA 9:7)

1. Is Fakul'tetskoy terapevticheskoy kliniki (dir. - prof. L.A. Varshamov) Saratovskogo meditsinskogo instituta.

(SLEEP, therapeutic use. hypertension, eff. on chronaximetry (Rus))

(HTFERTENSION, therapy, sleep ther., eff. on chronaximetry (Rus))

(MENUS? physiology, chronaximetry in hypertension after sleep ther. (Rus))
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BYREYEV, P.A., prof.; VAESHAMOV, L.A., prof.; VOLYNSKIY, B.G., dotsent; CERASIMOV, N.V., dotsent; CUREVICH, L.I., dotsent; ZHELYABOVSKIY, G.M., prof.; KARTASHOV, P.P., prof.; KOCHETOV, K.P., dotsent; KRUGLOV, A.N., prof.; KUTANIN, M.P., prof.; LARINA, V.S., dotsent; LOBKO, I.S., doktor [deceased]; LUKOVA, A.I., prof.; MAKHLIN, Ye.Yu., prof.; NAUMOV, A.I., kand.med.nauk; POPOV'YAN, I.M., prof.; SOLUN, H.S., kand.med.nauk; TARABUKHIN, M.M., dotsent; TRET'YAKOV, K.N., prof.; TRISHINA, A.A., kand.med.nauk; UL'YANOVA, A.V., dotsent; FAYN, A.E., kand.med.nauk; FAKTOROVICH, A.M., dotsent; FRANKFURT, A.I., prof.; FISHER, L.I., dotsent; CHASOVNIKOVA, Ye.P., kand.med.nauk; SHAMARIN, P.I., prof.; SHAPIRO, M.Ya., dotsent; SHVARTS, L.S., prof.; SHUSTERMAN, I.B., dotsent; FOY, A.M., prof.; FREYDMAN, S.L., kand.med.nauk; NIKITIN, B.A., dotsent, red.; AFANAS'YEV, I.A., red.; LUKASHEVICH, V., tekhn.red.

[Concise medical reference book] Kratkii terapevticheskii spravochnik. Izd.3., ispr. i dop. Saratov, Saratovskoe knizhnoe izd-vo, 1959. 919 p. (MIRA 13:7)

1. Chlen-korrespondent AMN SSSR (for Tret'yakov).

(MMDICINE-HANDBOOKS, MANUALS, ETC.)